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(54) Alkaline proteases

(57)Provided are an alkaline protease wherein an amino acid residue at (a) position 84, (b) position 104, (c) position 256 or (d) position 369 of SEQ ID NO:1 or at a position corresponding thereto has been deleted or selected from: at position (a): an arginine residue, at position (b): a proline residue, at position (c): an alanine, serine, glutamine, valine, leucine, asparagine, glutamic acid or aspartic acid residue, and at position (d): an aspartic acid residue; an alkaline protease wherein an amino acid residue at (e) position 66 or 264, (f) position 57, each of 101 to 106, 136, 193 or 342, (g) position 46 or 205, (h) position 54, 119, 138, 148 or 195, (i) position 247, (j) position 124, (k) position 107 or (l) position 257 has been deleted or selected from: at position (e): a glutamine, aspartic acid or like residue, at position (f): a lysine, serine or like residue, at position (g): a tyrosine

or tryptophan residue, at position (h): a tryptophan, phenylalanine or like residue, at position (i): a tryptophan, phenylalanine or like residue, at position (j): an alanine or lysine residue, at position (k): a lysine, arginine or like residue, and at position (l): a valine or isoleucine residue; a gene encoding the alkaline protease; a recombinant vector containing the gene, a transformant containing the recombinant vector; and a detergent composition containing the alkaline protease.

The present invention makes it possible to provide alkaline proteases having activity even in the presence of a high concentration of fatty acids, having high specific activity and detergency and being useful as an enzyme to be incorporated in a detergent.

Description

[0001] The present invention relates to alkaline proteases having high specific activity and strong oxidant resistance and as an enzyme to be added to a detergent, having excellent detergency.

[0002] Proteases have conventionally been used in a variety of fields such as various detergents including laundry detergents, cosmetics, bath agents, food modifiers, and pharmaceuticals such as digestion aids and anti-inflammatory agents. Among them, proteases for detergents are industrially produced in the largest amount and have a great market scale. A number of proteases are now put on the market.

[0003] In most cases, stains on clothes contain not only proteins but also plural components such as lipids and solid particles. There is accordingly a demand for detergents having detergency high enough to remove such actual complex stains. Finding, from such a viewpoint, alkaline proteases capable of retaining caseinolytic activity even in the presence of a high concentration of fatty acids and exhibiting excellent detergency even if the stain is composed of not a simple protein component but plural components, for example, protein and lipid, and having a molecular weight of about 43,000, the present inventors applied a patent (WO99/18218) on them.

[0004] Alkaline proteases superior to the above-described ones in specific activity, oxidant resistance and detergency and usable for detergents of wide-ranging compositions have however been requested.

[0005] The present inventors searched for such alkaline proteases mainly from enzyme variants. The above-described alkaline proteases are however utterly different in enzymological properties from serine proteases typified by subtilisin so that the modified site of subtilisin did not provide them with useful information. As a result of a further investigation, the present inventors have found that in order to obtain novel alkaline proteases having improved specific activity, stability against an oxidant and detergency while maintaining the properties of the above-described alkaline proteases, they must have a specific amino acid residue at a predetermined position of their amino acid sequence.

[0006] In one aspect of the present invention, there is thus provided an alkaline protease wherein an amino acid residue at (a) position 84, (b) position 104, (c) position 256 or (d) position 369 of SEQ ID NO:1 or at a position corresponding thereto has been deleted or selected from:

at position (a): an arginine residue,

at position (b): a proline residue,

at position (c): an alanine, serine, glutamine, valine, leucine, asparagine, glutamic acid or aspartic acid residue, and at position (d): an aspartic acid residue.

[0007] In another aspect of the present invention, there is also provided an alkaline protease wherein an amino acid residue at (e) position 66 or 264, (f) position 57, each of 101 to 106, 136, 193 or 342, (g) position 46 or 205, (h) position 54, 119, 138, 148 or 195, (i) position 247, (j) position 124, (k) position 107 or (l) position 257 of SEQ ID NO:1, or at a position corresponding thereto has been deleted or selected from:

at position (e): a glutamine, aspartic acid, serine, glutamic acid, alanine, threonine, leucine, methionine, cysteine, valine, glycine or isoleucine residue

at position (f): a lysine, serine, glutamine, phenylalanine, valine, arginine, tyrosine, leucine, isoleucine, threonine, methionine, cysteine, tryptophan, aspartic acid, glutamic acid, histidine, proline or alanine residue,

at position (g): a tyrosine, tryptophan, alanine, asparagine, glutamic acid, threonine, valine, leucine, isoleucine, histidine, serine, lysine, glutamine, methionine or cysteine residue,

at position (h): a tryptophan, phenylalanine, alanine, asparagine, glutamic acid, threonine, valine, histidine, serine, lysine, glutamine, methionine, glycine, aspartic acid, proline, arginine or cysteine residue,

at position (i): a tryptophan, phenylalanine, alanine, asparagine, glutamic acid, threonine, valine, leucine, isoleucine, histidine, serine, glutamine, methionine or cysteine residue,

at position (j): an alanine or lysine residue,

at position (k): a lysine, arginine, alanine or serine residue, and

at position (I): a valine or isoleucine residue.

[0008] In a further aspect of the present invention, there are also provided a gene encoding the alkaline protease, a recombinant vector containing the gene and a transformant containing the vector.

[0009] In a still further aspect of the present invention, there is also provided a detergent composition containing the alkaline protease.

FIG. 1 is a diagram illustrating detergency of an alkaline protease variant;

FIG. 2 is a diagram illustrating relative specific activity of each of alkaline protease variants; and FIG. 3 is a diagram illustrating relative residual activity of each of alkaline protease variants after treatment with an oxidant.

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[0010] As described above, in the alkaline proteases of the present invention, an amino acid residue at (a) position 84. (b) position 104, (c) position 256 or (d) position 369 of SEQ ID NO:1 or at a position corresponding thereto has been deleted or selected from: at position (a): an arginine residue, at position (b): a proline residue, at position (c): an alanine, serine, glutamine, valine, leucine, asparagine, glutamic acid or aspartic acid residue, and at position (d): an aspartic acid residue; or an amino acid residue at (e) position 66 or 264, (f) position 57, each of 101 to 106, 136, 193 or 342, (g) position 46 or 205, (h) position 54, 119, 138, 148 or 195, (i) position 247, (j) position 124, (k) position 107 or (I) position 257 of SEQ ID NO:1 or at a position corresponding thereto has been deleted or selected from: at position (e): a glutamine, aspartic acid, serine, glutamic acid, alanine, threonine, leucine, methionine, cysteine, valine, glycine or isoleucine residue, at position (f): a lysine, serine, glutamine, phenylalanine, valine, arginine, tyrosine, leucine, isoleucine, threonine, methionine, cysteine, tryptophan, aspartic acid, glutamic acid, histidine, proline or alanine residue, at position (g): a tyrosine, tryptophan, alanine, asparagine, glutamic acid, threonine, valine, leucine, isoleucine, histidine, serine, lysine, glutamine, methionine or cysteine residue, at position (h): a tryptophan, phenylalanine, alanine, asparagine, glutamic acid, threonine, valine, histidine, serine, lysine, glutamine, methionine, glycine, aspartic acid, proline, arginine or cysteine residue, at position (i): a tryptophan, phenylalanine, alanine, asparagine, glutamic acid, threonine, valine, leucine, isoleucine, histidine, serine, glutamine, methionine or cysteine residue, at position (j): an alanine or lysine residue, at position (k): a lysine, arginine, alanine or serine residue, and at position (l): a valine or isoleucine residue.

[0011] Specifically, the alkaline proteases according to the present invention mean alkaline proteases having an amino acid sequence represented by SEQ ID NO:1 wherein the amino acid residue at a position selected from the above-described (a) to (d) and (e) to (l) has been deleted or predetermined, or another alkaline protease wherein the amino acid residue at a position corresponding thereto has been deleted or predetermined. They may be wild type enzymes, wild type variants or artificial variants.

[0012] The "another alkaline protease" may be a wild type enzyme or a wild type variant. That having oxidant resistance and having a molecular weight, as determined by SDS-PAGE, of $43,000 \pm C2,000$ is preferred, of which that having an amino acid sequence showing at least 60% homology with the amino acid sequence of SEQ ID NO:1 is more preferred. Particularly preferred is that having an amino acid sequence showing at least 60% homology with the amino acid sequence of SEQ ID NO:1, having oxidant resistance, works on the alkaline side (pH 8 or greater), is stable with at least 80% residual activity when treated at pH 10 for 10 minutes even at 50°C, is inhibited by diisopropyl fluorophosphate (DFP) and phenylmethane sulfonyl fluoride (PMSF) and has a molecular weight, as determined by SDS-PAGE, of $43,000 \pm 2,000$. The term "having oxidant resistance" as used herein means that it has at least 50% of residual activity (synthetic substrate assay) when treated in a 50 mM hydrogen peroxide solution (containing 5 mM calcium chloride) at pH 10 (a 20 mM Britton-Robinson buffer) at 20°C for 20 minutes.

[0013] Examples of the "alkaline protease having an amino acid sequence represented by SEQ ID NO:1" include KP43 [derived from *Bacillus* sp. strain KSM-KP43 (FERM BP-6532), WO99/18218], while those of the "alkaline protease having an amino acid sequence showing at least 60% homology with the amino acid sequence of SEQ ID NO:1" include protease KP9860 having an amino acid sequence represented by SEQ ID NO:2 [derived from *Bacillus* sp. strain KSM-KP9860 (FERM BP-6534), WO99/18218], Protease E-1 having an amino acid sequence represented by SEQ ID NO:3 [derived from *Bacillus* sp. strain No. D-6 (FERM P-1592), JP740710], Protease Ya having an amino acid sequence represented by SEQ ID NO:4 [derived from *Bacillus* sp. strain Y (FERM BP-1029), JP861210], Protease SD521 having an amino acid sequence represented by SEQ ID NO:5 [derived from *Bacillus* sp. strain SD-521 (FERM BP-11162), JP910821], Protease A-1 having an amino acid sequence represented by SEQ ID NO:6 (derived from NCIB12289, WO8801293), and Protease A-2 having an amino acid sequence represented by SEQ ID NO:7 (derived from NCIB12513, WO8801293). Of these, the amino acid sequences selected from SEQ ID NOS. 2 to 7 or alkaline proteases showing at least 80%, more preferably at least 90%, especially at least 95% homology therewith are preferred.

[0014] The homology of an amino acid sequence is calculated by Lipman-Pearson's method (Science, 227, 1435 (1985)).

[0015] The "amino acid residue at a corresponding position" can be identified by comparing amino acid sequences by using known algorithm, for example, that of Lipman-Pearson. The position of the "amino acid residue at a corresponding position" in the sequence of each protease can be determined by aligning the amino acid sequence of the protease in such a manner. It is presumed that the corresponding position exists at the three-dimensionally same position in the amino acid sequence of SEQ ID NO:1 and the amino acid residue existing at the same position brings about similar effects for a specific function of the protease.

[0016] Described specifically,

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- (a) the amino acid residue at position 84 of SEQ ID NO:1 is a lysine residue. By employing the above-described method, the amino acid residue at the position corresponding thereto can be identified as the lysine residue at position 83 of SEQ ID NO:3. This amino acid residue is preferably arginine.
- (b) Although the amino acid residue at position 104 of SEQ ID NO:1 is a leucine residue, this amino acid residue

or an amino acid residue corresponding thereto is preferably a proline residue.

- (c) Although the amino acid residue at position 256 of SEQ ID NO:1 is a methionine residue, particularly preferred as this amino acid residue is an alanine, serine, glutamine, valine, leucine, asparagine, glutamic acid or aspartic acid residue.
- (d) Although the amino acid residue at position 369 of SEQ ID NO:1 is an aspartic acid residue, this amino acid residue or amino acid residue corresponding thereto is preferably an asparagine residue.
- (e) Although the amino acid residue at position 66 or 264 of SEQ ID NO:1 is an asparagine residue, this amino acid residue is preferably a glutamine, aspartic acid, serine, glutamic acid, alanine, threonine, leucine, methionine, cysteine, valine, glycine or isoleucine residue, with an aspartic acid, serine or glutamic acid residue being particularly preferred. More preferred is the case wherein the amino acid residue at position 66 is an aspartic acid residue and that at position 264 is a serine residue.
- (f) Although the amino acid residue at each of positions 57, 101 to 106, 136, 193 and 342 of SEQ ID NO:1 is a glycine residue, this amino acid residue is preferably a lysine, serine, glutamine, phenylalanine, valine, arginine, tyrosine, leucine, isoleucine, threonine, methionine, cysteine, tryptophan, aspartic acid, glutamic acid, histidine, proline or alanine residue. Particularly preferred is the case wherein the amino acid residue at position 57, 136, 193 or 342 is an alanine residue, or that at position 103 is an arginine residue.
- (g) Although the amino acid residue at position 46 or 205 of SEQ ID NO:1 is a phenylalanine residue, this amino acid residue is preferably a tyrosine, tryptophan, alanine, asparagine, glutamic acid, threonine, valine, leucine, isoleucine, histidine, serine, lysine, glutamine, methionine or cysteine residue. Particularly preferred is the case wherein the amino acid residue at position 46 is a leucine residue.
- (h) Although the amino acid residue at position 54, 119, 138, 148 or 195 of SEQ ID NO:1 is a tyrosine residue, this amino acid residue is preferably a tryptophan, phenylalanine, alanine, asparagine, glutamic acid, threonine, valine, histidine, serine, glutamine, methionine, glycine, aspartic acid, proline, lysine, arginine or cysteine residue. Particularly preferred is the case wherein the amino acid residue at position 195 is an alanine, aspartic acid, glutamic acid, glutamine, valine, tryptophan, glycine, lysine, threonine, methionine, cysteine, phenylalanine, proline, serine, arginine, asparagine or histidine residue.
- (i) Although the amino acid residue at position 247 of SEQ ID NO:1 is a lysine residue, this amino acid residue is preferably a tryptophan, phenylalanine, alanine, asparagine, glutamic acid, threonine, valine, leucine, isoleucine, histidine, serine, glutamine, methionine or cysteine residue. As the amino acid residue at position 247, an arginine or threonine residue is particularly preferred.
- (j) Although the amino acid residue at position 124 of SEQ ID NO:1 is an arginine residue, this amino acid residue is preferably an alanine or lysine residue.
- (k) Although the amino acid residue at position 107 of SEQ ID NO:1 is a leucine residue, this amino acid residue is preferably a lysine, arginine, alanine or serine residue, with a lysine residue being particularly preferred.
- (I) Although the amino acid residue at position 257 of SEQ ID NO:1 is an alanine residue, this amino acid residue is preferably a valine or isoleucine residue, with a valine residue being particularly preferred.

[0017] With regards to "another alkali protease" which is preferred among the above-exemplified ones, positions corresponding to (a) to (d) and (e) to (l) of the amino acid sequence (SEQ ID NO:1) of Protease KP43 and specific examples of an amino acid residue are shown below (Table 1-a, Table 1-b).

Table 1-a

				Proteases			
Position	TS43 SEQ ID NO:1	9860 SEQ ID NO:2	E-1 SEQ ID NO:3	Ya SEQ ID NO:4	SD-521 SEQ ID NO: 5	A-1 SEQ ID NO:6	A-2 SEQ ID NO:7
(a)	84Lys	84Lys	83Lys	83Lys	83Lys	84Lys	83Lys
(b)	104Leu	104Leu	103Leu	103Leu	103Leu	104Leu	103Leu
(c)	256Met	256Met	255Met	255Met	255Met	256Met	255Met
(d)	369Asp	369Asp	368Asp	368Asp	368Asp	369Asp	368Asp

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Table 1-b

					Proteases]
5	Position	TS43 SEQ ID NO:1	9860 SEQ ID NO:2	E-1 SEQ ID NO:3	Ya SEQ ID NO:4	SD-521 SEQ ID NO: 5	A-1 SEQ ID NO:6	A-2 SEQ ID NO:7
10	(e)	66Asn 264Asn	66Asn 264Asn	66Asn 263Asn	66Asn 263Asn	66Asn 263Asn	66Asn 264Asn	66Asn 263Asn
10	(f)	57Gly	57Gly	56Gly	56Gly	56Gly	57Gly	56Gly
		101Gly	101Ser	100Ser	100Ser	100Ser	101Asn	100Gly
		102Gly	102Gly	101Gly	101Gly	101Gly	102Gly	101Gly
15		103Gly	103Gly	102Gly	102Gly	102Gly	103Gly	102Gly
		105Gly	105Gly	104Gly	104Gly	104Gly	105Gly	104Gly
		106Gly	106Gly	105Gly	105Gly	105Gly	106Gly	105Gly
20		136Gly	136Gly	135Gly	135Gly	135Gly	136Gly	135Gly
20		193Gly	193Gly	192Gly	192Gly	192Gly	193Gly	192Gly
		342Gly	342Gly	341Gly	341Gly	341Gly	342Gly	341Gly
	(g)	46Phe	46Phe	46Phe	46Phe	46Phe	46Phe	46Phe
25		205Phe	205Phe	204Phe	204Phe	204Phe	205Phe	204Phe
	(h)	195Tyr	195Tyr	194lle	194IIe	194Leu	195Tyr	194Tyr
	(i)	247Lys	247Lys	246Lys	246Lys	246Lys	247Lys	246Lys
30	(j)	124Arg	124Arg	123Arg	123Arg	123Arg	124Arg	123Arg
	(k)	107Leu	107Leu	106Leu	106Leu	106Leu	107Leu	106Leu
	(I)	257Ala	257Ala	256Ala	256Ala	256Ala	257Ala	256Ala

[0018] In the alkaline proteases of the present invention, deletion of an amino acid residue or selection in (a) to (d) or (e) to (l) may be conducted at two or more positions simultaneously.

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[0019] When the alkaline protease of the present invention is a variant, the "protease having an amino acid sequence represented by SEQ ID NO:1" or the above-exemplified "another alkaline protease" serves as an alkaline protease prior to mutation (which may be called "parent alkaline protease"). By introducing mutation to a desired site of this parent alkaline protease, the alkaline protease of the present invention is available. For example, it is available by deleting or substituting, with another amino acid residue, the amino acid residue at a position selected from the above-described (a) to (d) and (e) to (l) of the amino acid sequence of SEQ ID NO:1 of Protease KP43 or at the corresponding position of the amino acid sequence of another alkaline protease, more specifically, amino acid sequence represented by SEQ ID NOS:2 to 7.

[0020] The alkaline protease of the present invention can be obtained, for example, by introducing mutation to a cloned gene encoding a parent alkaline protease, transforming a proper host by using the resulting mutated gene and then culturing the recombinant host. Cloning of the gene for encoding a parent alkaline protease may be carried out using an ordinary gene recombination technique, for example, in accordance with the process as described in WO99/18218, JP901128 or WO98/56927.

[0021] For mutation of a gene encoding a parent alkaline protease, either one of random mutation or site-specific mutation which is prevalent now can be adopted. More specifically, mutation can be effected using, for example, "Site-Directed Mutagenesis System Mutan-Super Express Kit" of Takara Shuzo Co., Ltd. By using recombinant PCR (polymerase chain reaction) as described in "PCR protocols" (Academic Press, New York, 1990), a desired sequence of a gene can be replaced with a sequence of another gene corresponding to the desired sequence.

[0022] For production of the protease variant of the present invention by using the resulting mutated gene, usable is, for example, the following process. A DNA encoding the protease variant of the present invention is stably amplified by linking the mutated gene with a DNA vector capable of amplifying it stably or by introducing the mutated gene onto a chromosomal DNA capable of maintaining it stably and then, the gene is introduced into a host permitting stable and

efficient expression of the gene, whereby the variant protease is produced. Hosts satisfying the above-described conditions include microorganisms belonging to *Bacillus* sp., *Escherichia coli*, mold, yeast and *Actinomyces*.

[0023] The alkaline protease of the present invention thus obtained has stable protease activity in an alkaline region, is free from the inhibition of caseinolytic activity by higher fatty acids, and has a molecular weight, as determined by SDS-PAGE, $43,000 \pm 2,000$. For example, the protease variant available from, as a parent strain, the protease having an amino acid sequence of SEQ ID NO:1 has the below-described physicochemical properties.

- (i) Acting pH range
- 10 [0024] It acts in a wide pH range of from 4 to 13 and exhibits at least 80% of the optimum pH active value at pH 6 to 12.
 - (ii) Stable pH range

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[0025] It is stable within a pH range of 6 to 11 under the treating conditions at 40°C for 30 minutes.

(iii) Influence of fatty acids

[0026] Its caseinolytic activity is not inhibited by oleic acid.

[0027] Such proteases of the present invention have excellent specific activity, oxidant resistance and detergency and are therefore useful as an enzyme to be incorporated in various detergent compositions. Particularly, the proteases wherein the amino acid residue at position (a) to (d) of SEQ ID NO:1 or at a position corresponding thereto has been deleted or specified are superior in detergency. Among them, those having, as the amino acid residue at (c) position 256 or at a position corresponding thereto, an alanine, serine, glutamine, valine, leucine, asparagine, glutamic acid or aspartic acid residue have both high specific activity and strong oxidant resistance. The proteases wherein the amino acid residue at position (e) to (I) of SEQ ID NO:1 or at a position corresponding thereto has been deleted or specified have particularly excellent specific activity.

[0028] The above-described protease may be added to the detergent composition of the present invention in an amount sufficient to permit exhibition of its activity. Although 0.1 to 5000 P.U. can be added per 1 kg of the detergent composition, 1000 P.U. or less, preferably 500 P.U. is added in consideration of economy.

[0029] To the detergent composition of the present invention, various enzymes can be used in combination with the alkaline protease of the present invention. Examples include hydrolases, oxidases, reductases, transferases, lyases, isomerases, ligases and synthetases. Of these, proteases, cellulases, lipases, keratinases, esterases, cutinases, amylases, pullulanases, pectinases, mannases, glucosidases, glucanases, cholesterol oxidases, peroxidases, laccases and proteases other than the alkaline protease used in the present invention are preferred.

[0030] Proteases include commercially available Alcalase, Esperase, Savinase and Everlase (each, product of Novo Nordisk), Properase and Purafect (each, product of Genencor International Inc.), and KAP (Kao Corp.) Cellulases include Cellzyme and Carezyme (each, product of Novo Nordisk), KAC (Kao Corp.) and alkaline cellulase produced by *Bacillus* sp. strain KSM-S237 as described in Japanese Patent Application Laid-Open No. Hei 10-313859. Amylases include Termamyl and Duramyl (each, product of Novo Nordisk), Purastar (Genencor International Inc.), and KAM (Kao Corp.). Lipases include Lipolase and Liporase Ultra (each, product of Novo Nordisk). The above-exemplified enzyme may be incorporated in an amount of 0.001 to 10%, preferably 0.03 to 5%.

[0031] A surfactant may be incorporated in an amount of 0.5 to 60 wt.% (which will hereinafter be called "%", simply) in the detergent composition. To a powdery detergent composition and a liquid detergent composition, addition of 10 to 45% and 20 to 50% are preferred, respectively. When the detergent composition of the present invention is a bleaching detergent or automatic dishwasher detergent, the surfactant may usually be added in an amount of 1 to 10%, preferably 1 to 5%.

[0032] A divalent metal ion scavenger may be added in an amount of 0.01 to 50%, with 5 to 40% being preferred.

[0033] An alkali agent and inorganic salt may be added in an amount of 0.01 to 80%, preferably 1 to 40%.

[0034] An antisoil redeposition agent may be added in an amount of 0.001 to 10%, preferably 1 to 5%.

[0035] A bleaching agent (ex. hydrogen peroxide or percarbonate) is added preferably in an amount of 1 to 10%.

Upon use of the bleaching agent, 0.01 to 10% of a bleaching activator can be added.

[0036] As a fluorescent brightener, biphenyl type ones (such as "Tinopal CBS-X") and stilbene type ones (such as DM fluorescent dye) can be used. It is added preferably in an amount of 0.001 to 2%.

[0037] The detergent composition of the present invention can be prepared in a conventional manner by using the alkaline protease obtained by the above-described process and the above-described known detergent components in combination. The detergent form can be selected according to the using purpose. Examples include liquid, powder and granule.

[0038] When the alkaline protease of the present invention is added to a powdery detergent composition, it is pre-

ferred to prepare detergent particles in advance and then mix therein the alkaline protease granules in accordance with the process as described in Japanese Patent Application Laid-Open No. Sho 62-25790 in order to avoid the contact of workers or end users with the enzyme upon preparation or use of the detergent or to prevent heat-induced deactivation or decomposition of the enzyme.

[0039] The detergent composition of the present invention thus available is usable as a laundry detergent, bleaching detergent, automatic dishwasher detergent, pipe cleaner and artificial tooth cleaner. Use as a laundry detergent, bleaching detergent or automatic dishwasher detergent is particularly preferred.

Example 1

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[0040] Mutation was introduced at random into a protease structural gene of about 2.0 kb including a termination codon by the following manner. First, PCR was conducted using a primer capable of amplifying this 2.0kb. A PCR master mix contained 5 ng of a template DNA, 20 pmoL of a phosphorylated primer, 20 nmoL of each dNTP, 1 μ moL of Tris/HCI (pH 8.3), 5 μ moL of KCI, 0.15 μ moL of MgCl₂ and 2.5U TaqDNA polymerase, and its total amount was adjusted to 100 μ L. After modification of the template by allowing it to stand at 94°C for 5 minutes, PCR was performed for 30 cycles, each cycle consisting of treatment at 94°C for 1 min, at 55°C for 1 min and at 72°C for 1.5 min. The PCR product was purified by "PCR product purification Kit" (product of Boeringer Manheim), followed by elution in 100 μ L of sterile water. With 1 μ L of the eluate, second PCR was conducted under conditions similar to those of the first PCR except for the template DNA. After completion of the second PCR, the PCR product was purified in a similar manner to the first PCR, followed by elution in 100 μ L of sterile water.

[0041] The amplified DNA fragment was integrated in a vector by polymerase reaction using "LATaq" produced by Takara Shuzo Co., Ltd. Described specifically, after addition of $5\,\mu\text{L}$ of a buffer for LATaq (a 10-fold concentrate), $8\,\mu\text{L}$ of a dNTP solution and $0.5\,\mu\text{L}$ of LATaq DNA polymerase, and as a template, 20 ng of plasmid pHA64TS (having a protease structural gene linked with an expression vector pHA64) to $35\,\mu\text{L}$ of the purified eluate, the total amount was adjusted to $50\,\mu\text{L}$. PCR reaction of the resulting liquid was carried out for 30 cycles, each consisting of treatment at 94°C for 1 min, 55° C for 1 min and 72°C for 4 min. By the subsequent ethanol precipitation, the PCR product was collected. This PCR product had a shape of a plasmid having a nick at the 5' prime end of the primer. Ligase reaction by T4 ligase (product of Takara Shuzo Co., Ltd.) was conducted to link this nick portion.

[0042] By using 10 μ L of this ligase reaction mixture, transformation of the *Bacillus subtilis* strain ISW1214 was conduced, whereby about 4×10^5 transformants were obtained. The resulting transformants of the strain ISW1214 were cultured on a skin-milk-containing medium (containing 1% skim milk, 1% bactotrypton, 1% sodium chloride, 0.5% yeast extract, 1.5% agar and 7.5 μ g/ml of tetracycline) and halo formation, which was presumed to reflect the protease secretion amount, was observed.

35 Example 2: Purification of an enzyme

[0043] The protease active fraction was prepared in the following manner. The transformants obtained in Example 1 was cultured at 30°C for 60 hours on a medium A (3% polypeptone S (product of Nippon Pharmaceutical), 0.5% yeast extract, 1% fish meat extract (product of Wako Pure Chemical Industries, Ltd.), 0.15% dipotassium phosphate, 0.02% magnesium sulfate 7 hydrate, 4% maltose and 7.5 μ g/mL of tetracycline). The supernatant of the thus-obtained cultured medium was added with ammonium sulfate to give 90% saturation, whereby salting-out of protein was caused. The sample obtained by salting-out was dissolved in a 10 mM tris HCl buffer (pH 7.5) containing 2 mM of calcium chloride. The resulting solution was dialyzed overnight against the same buffer by using a dialysis membrane. The fraction in the dialysis membrane was applied to DEAE Bio-Gel A (product of Bio-Rad Laboratories) equilibrated with a 10 mM tris HCl buffer (pH 7.5) containing 2 mM calcium chloride to collect the protease active fraction not adsorbed to the ion-exchanger. This active fraction was applied further to "SP-Toyopearl 550W" (product of Tosoh Corp.) equilibrated with the same buffer, followed by elution with a 0 to 50 mM sodium chloride solution, whereby a protease active fraction was obtained. The resulting fraction was analyzed by SDS-PAGE electrophoresis to confirm that the protease was obtained as substantially uniform protein. The protein concentration was measured in accordance with the method of Lowry, et al. (J. Biol. Chem. 193, 265-275(1981)) by using bovine serum albumin (product of Bio-Rad Laboratories) as a standard.

Example 3: Measuring method of protease activity

(1) Synthetic substrate assay

[0044] A decomposition rate was measured using a synthetic peptide made of Glt-Ala-Ala-Pro-Leu(A-A-P-L) as a substrate. Described specifically, a 50 mM borate/KCl buffer (pH 10.5) containing each enzyme to be evaluated and

3 mM of Glt-A-A-P-L-pNA (product of Peptide Institute, Inc) was kept at 30°C for 10 minutes and then, an absorbance at 420 nm was periodically measured. The peptide hydrolyzing activity was determined from an increasing ratio of the absorbance at 420 nm per unit hour. The protein was determined using a protein assay kit of Bio-Rad Laboratories.

5 (2) Natural substrate assay

[0045] After 1.0 mL of a 50 mM borate buffer (pH 10) containing 1% (w/v) of casein was kept at 30°C for 5 minutes, 0.1 mL of an enzyme solution was added and reaction was conducted for 15 minutes. To the reaction mixture, 2.0 mL of a reaction-stopping solution (0.11M trichloroacetic acid - 0.22M sodium acetate - 0.33M acetic acid) was added. The resulting mixture was allowed to stand at room temperature for 30 minutes and the filtered. The acid soluble protein in the filtrate was quantitatively determined by the modified method of Lowry, et al. Described specifically, after addition of 2.5 mL of an alkaline copper solution [1% potassium sodium tartrate : 1% copper sulfate : 1% sodium carbonate = 1:1:100] to the filtrate, the resulting solution was allowed to stand at room temperature for 10 minutes. Then, 0.25 mL of a diluted phenol solution (a phenol reagent (product of Kanto Kagaku) diluted 2-fold with ion exchange water) was added. After the resulting mixture was kept at 30°C for 30 minutes, absorbance at 660 nm was measured. One enzyme unit was designated as a quantity of the enzyme for liberating the acid soluble protein hydrolysis product corresponding to 1 mmol of tyrosine for 1 min in the above-described reaction.

Example 4

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(1) Preparation of granular detergent

[0046] Detergency of the detergent as described in Example 3 of WO99/29830 was evaluated. Described specifically, 465 kg of water was poured in a mixing tank of 1 m³ equipped with a stirring blade. After its water temperature reached 55°C, 48 kg of a 50% (w/v) aqueous solution of sodium dodecylenzenesulfonate and 135 kg of a 40% (w/v) aqueous solution of sodium polyacrylate were added. After stirring for 15 minutes, 120 kg of sodium carbonate, 60 kg of sodium sulfate, 9 kg of sodium sulfite and 3 kg of a fluorescent dye were added. After stirring for further 15 minutes, 300 kg of zeolite was added. The mixture was stirred for 30 minutes to yield a uniform slurry (the slurry had a water content of 50 wt.%). By spraying this slurry from a pressure spraying nozzle disposed in the vicinity of the top of a spray drying tower, base granules were obtained (a high temperature gas was fed to the spray drying tower at 225°C from the tower bottom and discharged from the tower top at 105°C).

[0047] Then, 15 parts by weight of a nonionic surfactant, 15 parts by weight of a 50 wt.% aqueous solution of sodium dodecylbenzenesulfonate and 1 part by weight of polyethylene glycol were mixed under heating to 70°C, whereby a mixture was obtained. In a Loedige mixer (product of Matsuzaka Giken Co., Ltd., capacity: 20L, equipped with a jacket), 100 parts by weight of the base granules obtained above were charged and stirring by a main shaft (150 rpm) and chopper (4000 rpm) was started. Warm water of 75°C was caused to flow in the jacket at 10 L/min, the mixture was charged therein in 3 minutes, and then stirring was conducted for 5 minutes. The surface of the detergent particles were covered with 10 parts by weight of crystalline aluminosilicate, whereby the final product of the granular detergent was obtained.

[Raw materials used]

Aqueous solution of sodium dodecylbenzenesulfonate: "Neopelex F65" (product of Kao Corp.)

[0048] Nonionic surfactant: "Emulgen 108KM" (product of Kao) added with 8.5 moles, on average, of ethylene oxide Aqueous solution of sodium polyacrylate: having an average molecular weight of 10000 (prepared in accordance with the process as described in Example of Japanese Patent Publication No. Hei 2-24283)

Sodium carbonate: dense ash (product of Central Glass Co., Ltd.)

Zeolite: "Zeolite 4A" having an average particle size of 3.5 µm (product of Tosoh Corp)

Polyethylene glycol: "K-PEG6000" (average molecular weight of 8500, product of Kao Corp.)
Fluorescent dye: "Tinopai CBS-X" (product of Ciba Geigy)

(2) Preparation of granulated protease

[0049] From the alkaline proteases of the present invention and a purified preparation of a parent alkaline protease, granulated protease was prepared based on the process as described in Japanese Patent Application Laid-Open No. Sho 62-257990 (6P.U./g)

(3) Measurement of detergency

[0050] In 1L of an aqueous calcium solution (71.2 mg calcium carbonate/1L) adjusted to 20°C, 0.67 g of each of the detergent compositions as shown in Table 2 was dissolved. With the resulting solution, a test cloth ("EMPA117" -(prepared by Swiss Federal Laboratories for Materials Testing and Research, blood/milk/carbon) cut into a piece of 6 × 6 cm was washed using a Terg-O-tometer (product of Ueshima Seisakusho) at 20°C and 100 rpm for 10 minutes. After rinsing and drying, the brightness was measured using a spectrophotometer ("CM3500d", product of MINOLTA). A detergency was calculated based on the below-described equation. The results are shown in Table 2.

[0051] Measuring results of the detergency of the protease variants obtained in Example 1 are shown in FIG. 1. The alkaline protease variants of the present invention each exhibited superior detergency to wild type enzymes to which mutation had not been introduced.

			Table 2						•
				Inven	tion pro	oducts		Comparati	ve products
			1	2	3	4	5	1	2
Parts by weight	Granulated	K84R	0.5						
	alkaline	L104P		0.5					
	proteas es	M256S			0.5				
	of this	M256A				0.5			
	invention	D369N					0.5		
	Granulated parent protease	alkaline						0.5	
	Granular detergen	t		99) .5	•			100
Detergency (%)			38	38	36	36	34	31	23

Example 5

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[0052] Measuring results, in accordance with the synthetic substrate assay or natural substrate assay, of protease activity of the protease variants obtained in Example 1 (the proteases modified at 195-position and 256-position amino acid residues, respectively was measured by the latter assay, while the other proteases were measured by the former assay) are shown in FIG. 2. The alkaline protease variants of the present invention exhibited high specific activity.

Example 6

[0053] In 2 mL of a 100 mM borate buffer (pH 10.5) containing 3% of aqueous hydrogen peroxide, a 50 µl portion of each of the protease variants obtained by purification in Example 1 was added. The resulting mixture was allowed to stand at 30°C for 30 minutes. After addition of an adequate amount of catalase (product of Boehringer Manheim) to remove excess hydrogen peroxide, the residual protease activity was measured by the synthetic substrate assay. In FIG. 3, the residual activity after treatment with aqueous hydrogen peroxide is shown relative to the activity before treatment set at 100%.

[0054] The alkaline protease variants of the present invention exhibited higher oxidant resistance than the parent alkaline protease.

[0055] The present invention makes it possible to provide alkaline proteases having activity even under a high concentration of fatty acids, having high specific activity, oxidant resistance and detergency, and being useful as an enzyme to be incorporated in a detergent.

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15	Ser Tyr Pro Tyr 385	Asp Asn Asn Trp As 390	p Gly Arg Asn Asn Val 395	Glu Asn 400
	Val Phe Ile Asr	Ala Pro Gln Ser Gl 405	y Thr Tyr Thr Ile Glu 410	Val Gln 415
20	Ala Tyr Asn Val		n Arg Phe Ser Leu Ala 5 430	Ile Val
	His		-	
25	<210> 6 <211> 434 <212> PRT <213> Bacillus	sp.		
30	<400> 6 Asn Asp Val Ala 1	a Arg Gly Ile Val Ly 5	rs Ala Asp Val Ala Gln 10	Ser Ser 15
	Tyr Gly Leu Tyr	_	al Val Ala Val Ala Asp 25 30	
<i>35</i>	Leu Asp Thr Gly	y Arg Asn Asp Ser Se 40	er Met His Glu Ala Phe 45	Arg Gly
40	Lys Ile Thr Ala	a Ile Tyr Ala Leu Gl 55	y Arg Thr Asn Asn Ala 60	Asn Asp
	Pro Asn Gly Hi:	s Gly Thr His Val Al 70	a Gly Ser Val Leu Gly 75	Asn Gly 80
45	Thr Ser Asn Ly	s Gly Met Ala Pro Gl 85	in Ala Asn Leu Val Phe 90	Gln Ser 95
	Val Met Asp Se:		ly Gly Leu Pro Ser Asn 05 110	
50	Thr Leu Phe Sei	r Gln Ala Tyr Ser Al 120	la Gly Ala Arg Ile His 125	Thr Asn
<i>55</i>	Ser Trp Gly Al	a Pro Val Asn Gly Al 135	la Tyr Thr Thr Asp Ser 140	Arg Asn

	Val 145	Asp	Asp	Tyr	Val	Arg 150	Lys	Asn	Asp	Met	Ala 155	Val	Leu	Phe	Ala	Ala 160
5	Gly	Asn	Glu	Gly	Pro 165	Asn	Gly	Gly	Thr	Ile 170	Ser	Ala	Pro	Gly	Thr 175	Ala
10	Lys	Asn	Ala	Ile 180	Thr	Val	Gly	Ala	Thr 185	Glu	Asn	Leu	Arg	Pro 190	Ser	Phe
	Gly	Ser	Tyr 195	Ala	Asp	Asn	Ile	Asn 200	His	Val	Ala	Gln	Phe 205	Ser	Ser	Arg
15	Gly	Pro 210	Thr	Lys	Asp	Gly	Arg 215	Ile	Lys	Pro	Asp	Val 220	Met	Ala	Pro	Gly
	Thr 225	Phe	Ile	Leu	Ser	Ala 230	Arg	Ser	Ser	Leu	Ala 235	Pro	Asp	Ser	Ser	Phe 240
20	Trp	Ala	Asn	His	Asp 245	Ser	Lys	Tyr	Ala	Tyr 250	Met	Gly	Gly	Thr	Ser 255	Met
25	Ala	Thr	Pro	Ile 260	Val	Ala	Gly	Asn	Val 265	Ala	Gln	Leu	Arg	Glu 270	His	Phe
	Ile	Lys	Asn 275	Arg	Gly	Ile	Thr	Pro 280	Lys	Pro	Ser	Leu	Leu 285	Lys	Ala	Ala
30	Leu	Ile 290	Ala	Gly	Ala	Thr	Asp 295	Ile	Gly	Leu	Gly	Tyr 300	Pro	Ser	Gly	Asn
	Gln 305	Gly	Trp	Gly	Arg	Val 310	Thr	Leu	Asp	Lys	Ser 315	Leu	Asn	Val	Ala	Phe 320
35	Val	Asn	Glu	Thr	Ser 325	Ser	Leu	Ser	Thr	Asn 330	Gln	Lys	Ala	Thr	Tyr 335	Ser
	Phe	Thr	Ala	Gln 340	Ser	Gly	Lys	Pro	Leu 345	Lys	Ile	Ser	Leu	Val 350	Trp	Ser
40	Asp	Ala	Pro 355	Ala	Ser	Thr	Ser	Ala 360	Ser	Val	Thr	Leu	Val 365	Asn	Asp	Leu
45	Asp	Leu 370	Val	Ile	Thr	Ala	Pro 375	Asn	Gly	Thr	Lys	Туг 380	Val	Gly	Asn	Asp
	Phe 385	Thr	Ala	Pro	Tyr	Asp 390	Asn	Asn	Trp	Asp	Gly 395	Arg	Asn	Asn	Val	Glu 400
50	Asn	Val	Phe	Ile	Asn 405	Ala	Pro	Gln	Ser	Gly 410	Thr	Tyr	Thr	Val	Glu 415	Val
	Gln	Ala	Tyr	Asn 420	Val	Pro	Gln	Gly	Pro 425	Gln	Ala	Phe	Ser	Leu 430	Ala	Ile
55	Val	Asn														

5	<210> 7 <211> 433 <212> PRT <213> Bacillus sp.												
10	400> 7 sn Asp Val Ala Arg Gly Ile Val Lys Ala Asp Val Ala Gln Asn A 1 5 10 15	sn											
	the Gly Leu Tyr Gly Gln Gly Gln Ile Val Ala Val Ala Asp Thr G 20 25 30	ly											
15	eu Asp Thr Gly Arg Asn Asp Ser Ser Met His Glu Ala Phe-Arg G	ly											
	ys Ile Thr Ala Leu Tyr Ala Leu Gly Arg Thr Asn Asn Ala Asn A 50 55 60	qe											
20	Pro Asn Gly His Gly Thr His Val Ala Gly Ser Val Leu Gly Asn A	1a 80											
25	hr Asn Lys Gly Met Ala Pro Gln Ala Asn Leu Val Phe Gln Ser I 85 90 95	le											
25	Met Asp Ser Gly Gly Leu Gly Gly Leu Pro Ala Asn Leu Gln T 100 105 110	.hr											
30	eu Phe Ser Gln Ala Tyr Ser Ala Gly Ala Arg Ile His Thr Asn S 115 120 125	er											
	Trp Gly Ala Pro Val Asn Gly Ala Tyr Thr Thr Asp Ser Arg Asn V 130 135 140	/al											
35	asp Asp Tyr Val Arg Lys Asn Asp Met Thr Ile Leu Phe Ala Ala G 150 155 1	60 60											
	Asn Glu Gly Pro Gly Ser Gly Thr Ile Ser Ala Pro Gly Thr Ala L 165 170" 175	ıys											
40	Asn Ala Ile Thr Val Gly Ala Thr Glu Asn Leu Arg Pro Ser Phe G 180 185 190	ly											
45	Ser Tyr Ala Asp Asn Ile Asn His Val Ala Gln Phe Ser Ser Arg G 195 200 205	ly											
	Pro Thr Arg Asp Gly Arg Ile Lys Pro Asp Val Met Ala Pro Gly T 210 215 220	hr											
50	Cyr Ile Leu Ser Ala Arg Ser Ser Leu Ala Pro Asp Ser Ser Phe T 235 235 2	rp 240											
	Ala Asn His Asp Ser Lys Tyr Ala Tyr Met Gly Gly Thr Ser Met A 245 250 255	lla											
55													

	Thr	Pro	Ile	Val 260	Ala	Gly	Asn	Val	Ala 265	Gln	Leu	Arg	Glu	His 270	Phe	Val
5	Lys	Asn	Arg 275	Gly	Val	Thr	Pro	Lys 280	Pro	Ser	Leu	Leu	Lys 285	Ala	Ala	Leu
10	Ile	Ala 290	Gly	Ala	Ala	Asp	Val 295	Gly	Leu	Gly	Phe	Pro 300	Asn	Gly	Asn	Gln
	Gly 305	Trp	Gly	Arg	Val	Thr 310	Leu	Asp	Lys	Ser	Leu 315	Asn	Val	Ala	Phe	Val 320
15	Asn	Glu	Thr	Ser	Pro 325	Leu	Ser	Thr	Ser	Gln 330	Lys	Ala	Thr	_	Ser 335	Phe
	Thr	Ala	Gln	Ala 340	Gly	Lys	Pro	Leu	Lys 345	Ile	Ser	Leu	Val	Trp 350	Ser	Asp
20	Ala	Pro	Gly 355	Ser	Thr	Thr	Ala	Ser 360	Leu	Thr	Leu	Val	Asn 365	Asp	Leu	Asp
	Leu	Val 370	Ile	Thr	Ala	Pro	Asn 375	Gly	Thr	Lys	Tyr	Val 380	Gly	Asn	Asp	Phe
25	Thr 385	Ala	Pro	Tyr	Asp	Asn 390	Asn	Trp	Asp	Gly	Arg 395	Asn	Asn	Val	Glu	Asn 400
30	Val	Phe	Ile	Asn	Ala 405	Pro	Gln	Ser	Gly	Thr 410	Tyr	Thr	Val	Glu	Val 415	Gln
	Ala	Tyr	Asn	Val 420	Pro	Val	Ser	Pro	Gln 425	Thr	Phe	Ser	Leu	Ala 430	Ile	Val
<i>35</i>	His															

Claims

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1. An alkaline protease wherein an amino acid residue at (a) position 84, (b) position 104, (c) position 256 or (d) position 369 of SEQ ID NO:1 or at a position corresponding thereto has been deleted or selected from:

at position (a): an arginine residue,

at position (b): a proline residue,

at position (c): an alanine, serine, glutamine, valine, leucine, asparagine, glutamic acid or aspartic acid residue, and

at position (d): an aspartic acid residue.

2. An alkaline protease having an amino acid sequence represented by SEQ ID NO:1 or having an amino acid sequence showing at least 60% homology therewith, wherein an amino acid residue at (a) position 84, (b) position 104, (c) position 256 or (d) position 369 of SEQ ID NO:1 or at a position corresponding thereto has been deleted or selected from:

at position (a): an arginine residue,

at position (b): a proline residue,

at position (c): an alanine, serine, glutamine, valine, leucine, asparagine, glutamic acid or aspartic acid residue, and

at position (d): an aspartic acid residue.

3. An alkaline protease wherein an amino acid residue at (e) position 66 or 264, (f) position 57, each of 101 to 106, 136, 193 or 342, (g) position 46 or 205, (h) position 54, 119, 138, 148 or 195, (i) position 247, (j) position 124, (k) position 107 or (l) position 257 of SEQ ID NO:1, or at a position corresponding thereto has been deleted or selected from::

at position (e): a glutamine, aspartic acid, serine, glutamic acid, alanine, threonine, leucine, methionine, cysteine, valine, glycine or isoleucine residue

at position (f): a lysine, serine, glutamine, phenylalanine, valine, arginine, tyrosine, leucine, isoleucine, threonine, methionine, cysteine, tryptophan, aspartic acid, glutamic acid, histidine, proline or alanine residue, at position (g): a tyrosine, tryptophan, alanine, asparagine, glutamic acid, threonine, valine, leucine, isoleucine, histidine, serine, lysine, glutamine, methionine or cysteine residue,

at position (h): a tryptophan, phenylalanine, alanine, asparagine, glutamic acid, threonine, valine, histidine, serine, lysine, glutamine, methionine, glycine, aspartic acid, proline, arginine or cysteine residue,

at position (i): a tryptophan, phenylalanine, alanine, asparagine, glutamic acid, threonine, valine, leucine, isoleucine, histidine, serine, glutamine, methionine or cysteine residue,

at position (j): an alanine or lysine residue,

at position (k): a lysine, arginine, alanine or serine residue, and

at position (I): a valine or isoleucine residue.

4. An alkaline protease having an amino acid sequence represented by SEQ ID NO:1 or having an amino acid sequence showing at least 60% homology therewith, wherein an amino acid residue at (e) position 66 or 264, (f) position 57, each of 101 to 106, 136, 193 or 342, (g) position 46 or 205, (h) position 54, 119, 138, 148 or 195, (i) position 247, (j) position 124, (k) position 107 or (l) position 257 has been deleted or selected from:

at position (e): a glutamine, aspartic acid, serine, glutamic acid, alanine, threonine, leucine, methionine, cysteine, valine, glycine or isoleucine residue

at position (f): a lysine, serine, glutamine, phenylalanine, valine, arginine, tyrosine, leucine, isoleucine, threonine, methionine, cysteine, tryptophan, aspartic acid, glutamic acid, histidine, proline or alanine residue, at position (g): a tyrosine, tryptophan, alanine, asparagine, glutamic acid, threonine, valine, leucine, isoleucine, histidine, serine, lysine, glutamine, methionine or cysteine residue,

at position (h): a tryptophan, phenylalanine, alanine, asparagine, glutamic acid, threonine, valine, histidine, serine, lysine, glutamine, methionine, glycine, aspartic acid, proline, arginine or cysteine residue,

at position (i): a tryptophan, phenylalanine, alanine, asparagine, glutamic acid, threonine, valine, leucine, isoleucine, histidine, serine, glutamine, methionine or cysteine residue,

at position (j): an alanine or lysine residue,

at position (k): a lysine, arginine, alanine or serine residue, and

at position (I): a valine or isoleucine residue.

5. An alkaline protease according to claim 2 or 4, wherein the amino acid sequence represented by SEQ ID NO:1 or amino acid sequence showing at least 60% homology therewith is an amino acid sequence selected from SEQ ID NOS: 2 to 7.

- 6. A gene encoding an alkaline protease as claimed in any one of claims 1 to 5.
 - 7. A recombinant vector comprising a gene as claimed in claim 6.
 - 8. A transformant comprising a recombinant vector as claimed in claim 7.

9. A transformant according to claim 8, wherein a microorganism is used as a host.

10. A detergent composition comprising an alkaline protease as claimed in any one of claims 1 to 5.

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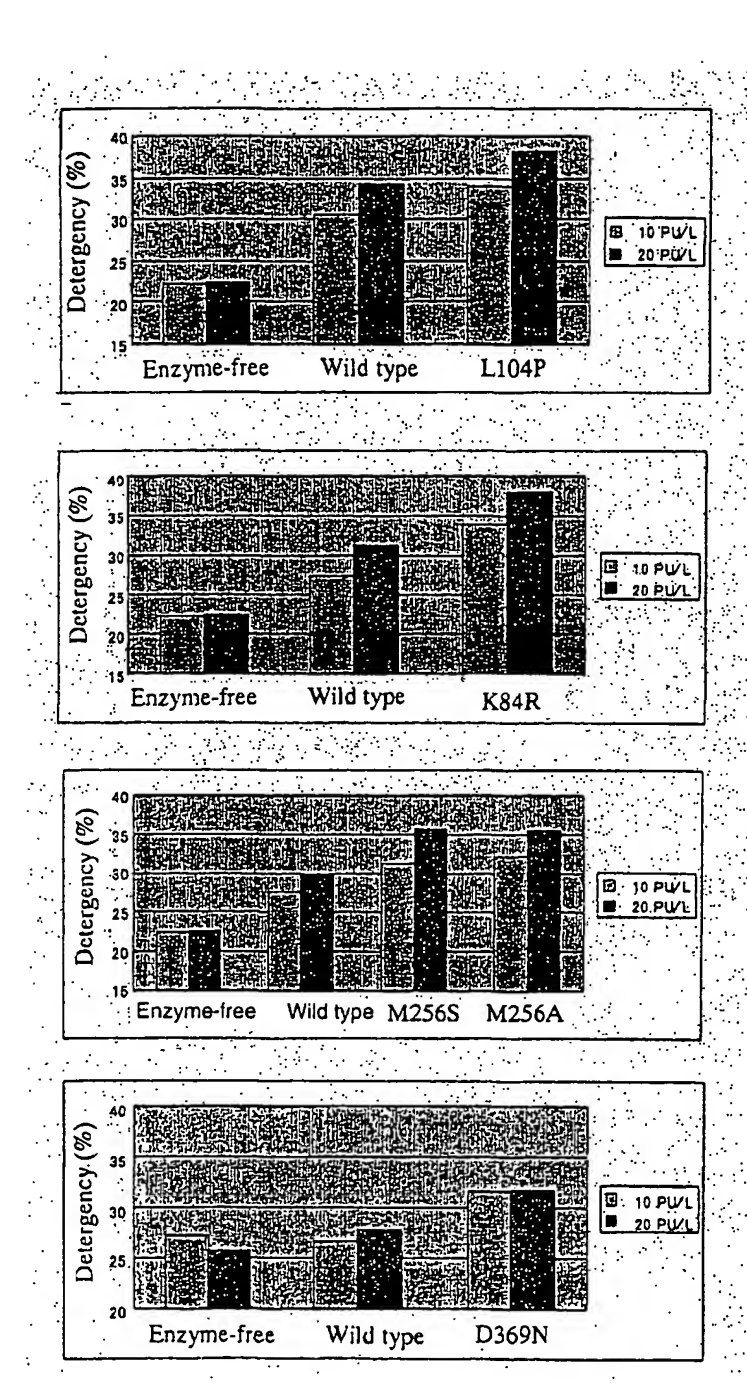
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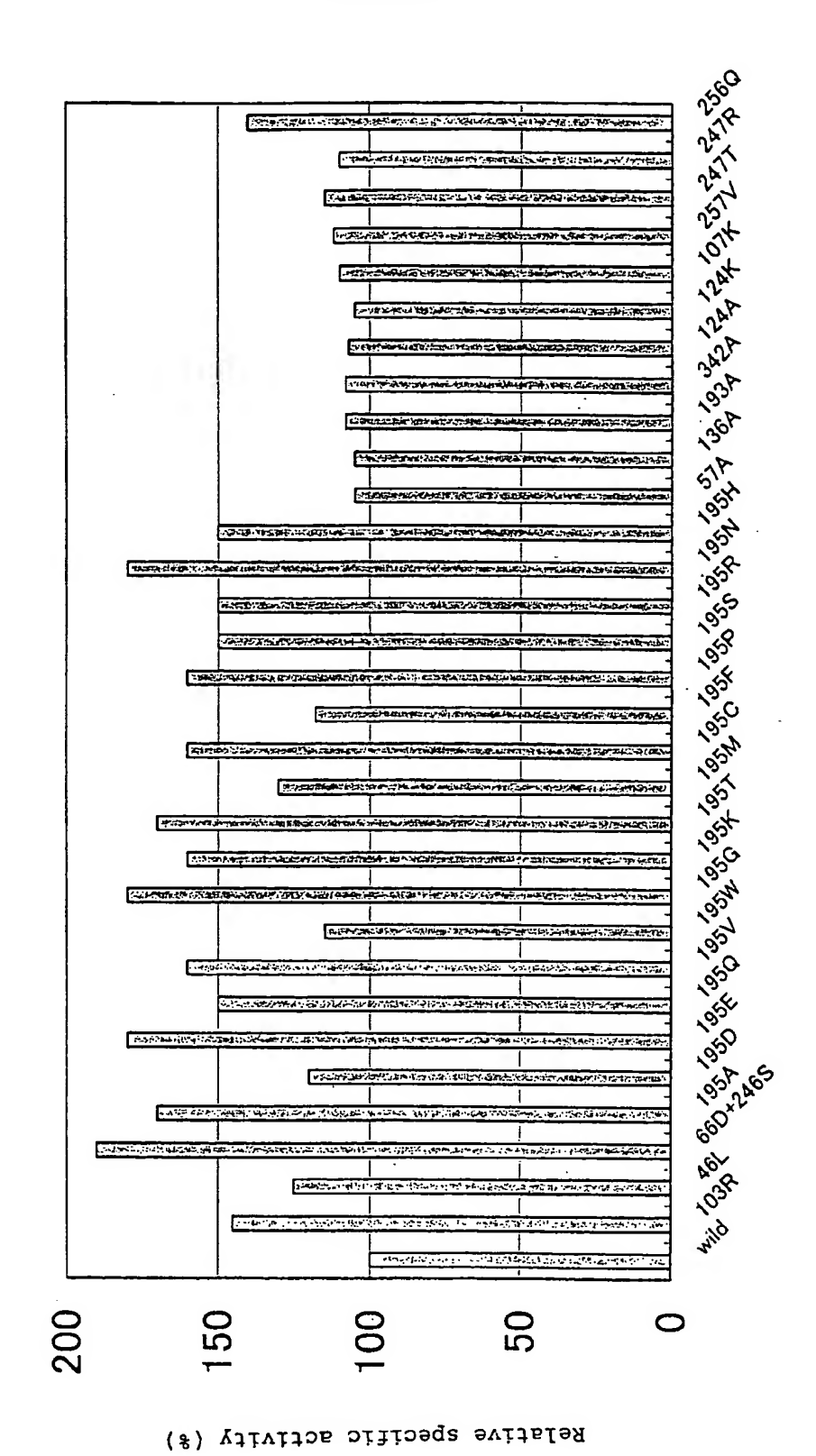
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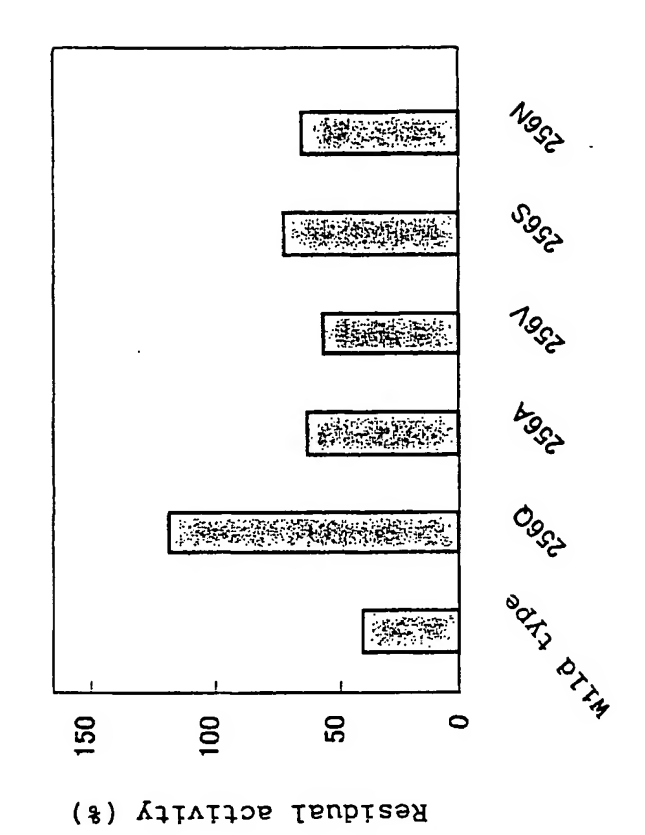
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FIG. 1





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